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INFORMATION DISCLOSURE STATEMENT BY APPLICANT <i>(Use several sheets if necessary)</i>		APPLICANT Allen et al.	
		FILING DATE 16 Jun 2006	GROUP


U.S. PATENT DOCUMENTS

Examiner Initial	Document Number	Date	Name	Class	Subclass	Filing Date If Appropriate
22	US-2006/0089375-A1	27 Apr 2006	Allen, Coe, Cook, et al.			
	US-3,755,340	28 Aug 73	Hoehn et al.			
	US-3,833,594	03 Sep 74	Hoehn et al. / E.R. Squibb & Sons, Inc.			
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22	US-3,979,399	07 Sep 76	Hoehn et al. / E.R. Squibb & Sons, Inc.			

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	Document Number	Date	Country	Class	Subclass	Translation	
						Yes	No
22	CA-1003419		CA				
22	WO-02/081463-A1	17 Oct 2002	WO				

OTHER DOCUMENTS (Including Author, Title, Date, Pertinent Pages, Etc.)

22	BARE T.M. ET AL.; Synthesis and structure-activity relationships of a series of anxiolytic pyrazolopyridine ester and amide anxiolytic agents; Journal of Medicinal Chemistry; 1989; 32; pages 2561-2573;
22	HOEHN H. ET AL.; 1H-pyrazolo[3,4-b]pyridines; Journal of Heterocyclic Chemistry; 1972; 9(2); pages 235-253;
22	OCHIAI H. ET AL.; New orally active PDE4 inhibitors with therapeutic potential; Bioorg. Med. Chem. Lett.; 5th Jan 2004 issue (available as "articles in press" version on or before 4th December 2003, possibly October 2003, via internet); 14(1); pages 29-32;
22	SCHENONE S. ET AL.; Synthesis and biological data of 4-amino-1-(2-chloro-2-phenylethyl)-1H-pyrazolo[3,4-b]pyridine-5-carboxylic acid ethyl esters, a new series of A1-adenosine receptor (A1AR) ligands; Bioorg. Med. Chem. Lett.; 2001; 11; pages 2529-2531;
22	YU G., MASON H.J., ET. AL.; Substituted pyrazolopyridines as potent and selective PDE5 inhibitors: potential agents for treatment of erectile dysfunction; Journal of Medicinal Chemistry; 2001; 44; pages 1025-1027;
EXAMINER 	
DATE CONSIDERED 10/17/07	
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